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LIST OF DOCUMENTARY INFORMATION CITED BY APPLICANT (Use several sheets if necessary)	APPLICANT	Lobel, et al.
	FILING DATE	May 9, 2001
	GROUP	1631 1632

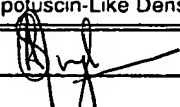
U.S. PATENT DOCUMENTS

		DOCUMENT NUMBER	DATE	NAME	CLASS	SUB- CLASS	FILING DATE IF APPROPRIATE

FOREIGN PATENT DOCUMENTS

		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUB- CLASS	TRANSLATION YES NO

OTHER PRIOR ART (including Author, Title, Date, Pertinent Pages, Etc.)

af	AA	Sharp, et al., Human Molecular Genetics, 1997, 6:591-595, Loci for classical and a variant late infantile neuronal ceroid lipofuscinosis map to chromosomes 11p15 and 15q21-23
af	AB	Lerner, et al., Cell, 1995, 82:949-957, Isolation of a Novel Gene Underlying Batten Disease, CLN3.
af	AC	Ivy, et al., American Journal of Medical Genetics, 1992, 42:555-560, Protease Inhibitors as a Model for NCL Disease, with Special Emphasis on the Infantile and Adult Forms.
af	AD	Ashizawa, et al., American Journal of Medical Genetics, 1992, 42:55-60, Diagnostic Value of Ophthalmologic Finding in Myotonic Dystrophy: Comparison with Risks Calculated by Haplotype Analysis of Closely Linked Restriction Fragment Length Polymorphisms.
af	AE	Sleat, et al., Biochem, 1997, 324:33-39, - Glucosidase and N-acetylglucosamine-6- sulphatase are the major mannose-6-phosphate glycoproteins in human urine.
af	AF	Ezaki, et al., Journal of Neurochemistry, 1996, 67:1677-1687, Specific Delay in the degradation of Mitochondrial ATP Synthase Subunit c in Late Infantile Neuronal Ceroid Lipofuscinosis is Derived from Cellular Proteolytic Dysfunction rather than Structural Alteration of Subunit c.
af	AG	Ivey, et al., Science, 1994, 226:985-987, Inhibitors of Lysosomal enzymes: Accumulation of Lipofuscin-Like Dense Bodies in the Brain
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